

Functional Electrospun Polystyrene Nanofibers Incorporating α -, β -, and γ -Cyclodextrins: Comparison of Molecular Filter Performance

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Electrospinning of functional polymeric nanofibers has attracted considerable attention in the past decade due to the simplicity of the process and the enhanced properties associated with the size of the fibers.^{1–4} One potential application for electrospun nanofibers is in the field of filtration and separation science where the so-called nanoweb can provide separation of tiny particles and have significantly higher adsorptive capacity in filtration.^{5–8} In filtration of organic molecules from complex solutions, the capacity to selectively remove the target molecule(s) with high efficiency is highly desirable. One approach that we employ is to exploit the properties of cyclodextrins (CDs) that can form inclusion complexes with hazardous chemicals and polluting substances.^{9–13} Hence, the functionalization of electrospun nanofibers/nanoweb with CDs incorporated is very appealing since these nanoweb can potentially act as highly selective molecular filters for the removal of different types of organic molecules from solution.^{14,15} In addition, the high surface area of the nanofibers significantly enhances the uptake efficiency as more CDs are present.

Cyclodextrins (CDs) are cyclic oligosaccharides having a toroid-shaped molecular structure that are able to form noncovalent host–guest complexes with various molecules.^{16–19} The most common CDs can have 6, 7, or 8 glucopyranose units in the cyclic and are referred to as α -, β -, and γ -cyclodextrins (α -CD, β -CD, γ -CD), respectively (Figure 1). The depth of the cavities for the three CDs is the same (~ 7.8 Å), while

ABSTRACT Electrospinning has been used to successfully create polystyrene (PS) nanofibers containing either of three different types of cyclodextrin (CD); α -CD, β -CD, and γ -CD. These three CDs are chosen because they have different sized cavities that potentially allow for selective inclusion complex (IC) formation with molecules of different size or differences in affinity of IC formation with one type of molecule. The CD containing electrospun PS nanofibers (PS/CD) were initially characterized by scanning electron microscopy (SEM) to determine the uniformity of the fibers and their fiber diameter distributions. X-ray photoelectron spectroscopy (XPS) was used to quantitatively determine the concentration of each CD on the different fiber surfaces. Static time-of-flight secondary ion mass spectrometry (*static*-ToF-SIMS) showed the presence of each type of CD on the PS nanofibers by the detection of both the CD sodium adduct molecular ions ($M + Na^+$) and lower molecular weight oxygen containing fragment ions. The comparative efficiency of the PS/CD nanofibers/nanoweb for removing phenolphthalein, a model organic compound, from solution was determined by UV–vis spectrometry, and the kinetics of phenolphthalein capture was shown to follow the trend $PS/\alpha\text{-CD} > PS/\beta\text{-CD} > PS/\gamma\text{-CD}$. Direct pyrolysis mass spectrometry (DP-MS) was also performed to ascertain the relative binding strengths of the phenolphthalein for the CD cavities, and the results showed the trend in the interaction strength was $\beta\text{-CD} > \gamma\text{-CD} > \alpha\text{-CD}$. Our results demonstrate that nanofibers produced by electrospinning that incorporate cyclodextrins with different sized cavities can indeed filter organic molecules and can potentially be used for filtration, purification, and/or separation processes.

KEYWORDS: cyclodextrin · electrospinning · polystyrene · nanofiber · molecular filter · nanoweb

the cavity diameters of α -CD, β -CD, and γ -CD are ~ 6 , 8, and 10 Å, respectively.²⁰ The formation of cyclodextrin inclusion complex (CD-IC) and its strength/stability mainly depend on the size/shape match and binding forces (hydrophobic interactions, van der Waals attractions, hydrogen bonding, electrostatic interactions, *etc.*) between the host CD cavity and guest molecules.^{21,22} Therefore, different types of CDs (α -CD, β -CD, γ -CDs, and chemically modified CD derivatives) show different capabilities and stabilities when forming inclusion complexes with the same host molecule.

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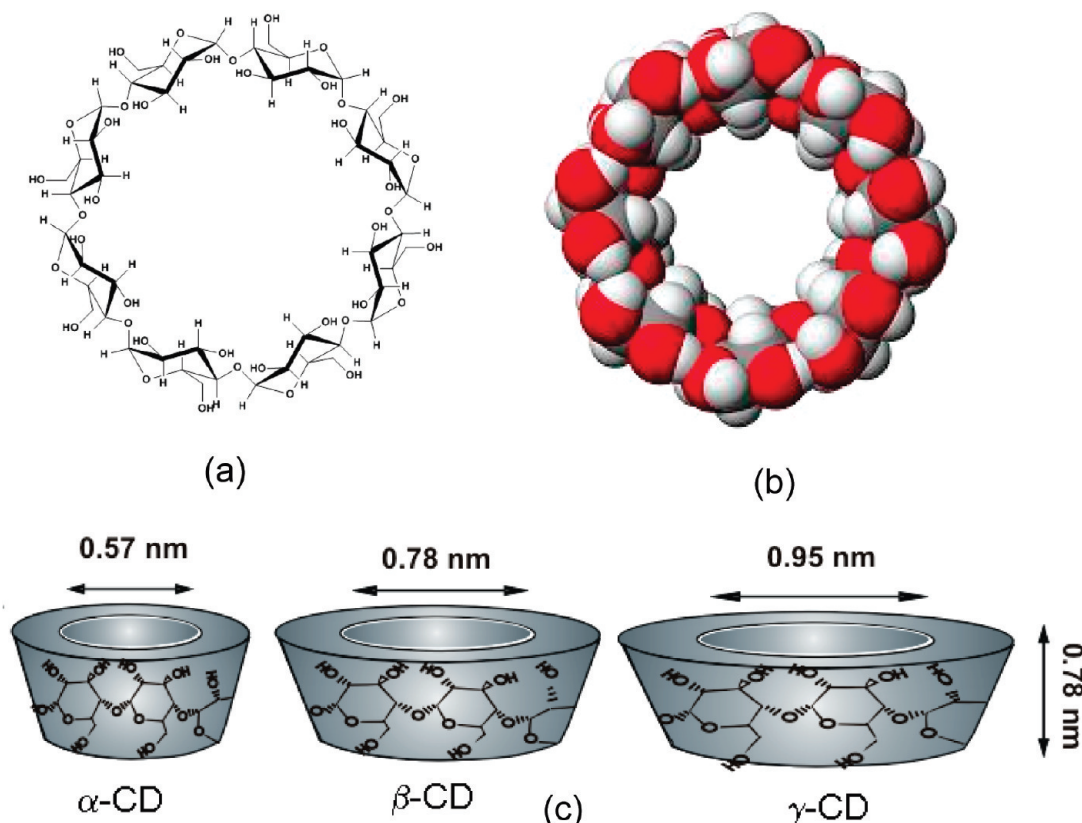


Figure 1. (a) Chemical structure of γ -CD, (b) 3-D structure of γ -CD, and (c) approximate dimensions of α -CD, β -CD, and γ -CD.²⁰

In our very recent studies, we have shown that functional electrospun nanofibers can be obtained by incorporating cyclodextrins (CDs) and cyclodextrin inclusion complexes (CD-ICs) into nanofibers by electrospinning.^{14,23–28} In one of these studies, we have shown that electrospun polystyrene (PS) fibers incorporating β -CD have the potential to be used as molecular filters.¹⁴ Here we extend on this work by electrospinning polystyrene (PS) with three types of CDs (α -CD, β -CD, and γ -CDs) separately, aimed at investigating their relative efficiency at filtering organic molecules from solution. Paramount to the success of the PS/CD nanofibers functioning as molecular filters is the presence of the CD molecules at the fiber surface. We have utilized X-ray photoelectron spectroscopy (XPS) and static time-of-flight secondary ion mass spectrometry (static-ToF-SIMS) to detect the CD molecules in the outer molecular layers of the fiber surfaces. The ability of the PS/CD nanofibers to filter molecules from solution was determined by using phenolphthalein as a

model compound, and the removal from solution was detected using UV–vis spectrophotometry. Finally, the relative affinity of the phenolphthalein for the different CD molecules was determined using direct pyrolysis mass spectrometry (DP-MS), and the results indicate that the cavities of the CDs have different binding strengths for phenolphthalein. Thus, electrospun nanofibers functionalized with different CD molecules may have tunable properties that can be exploited to manufacture molecular filters for highly selective and efficient removal of organic molecules from solution.

RESULTS AND DISCUSSION

SEM Characterization. The optimization of the electrospinning conditions for producing bead-free uniform PS fibers incorporating cyclodextrins (PS/ β -CD) and the bulk chemical characterization of these electrospun PS and PS/ β -CD fibers was reported elsewhere.²⁶ The composition and the properties of the PS and PS/CD solutions and the resulting electrospun fibers are summa-

TABLE 1. Properties of PS and PS/CD Solutions and the Average Fiber Diameter of the Resulting Electrospun Fibers

materials	% PS (w/v) ^a	CD type, % (w/w) ^b	viscosity (cP)	conductivity (μ S/cm)	fiber diameter range (nm)	average fiber diameter (nm)
PS	25		240.4 \pm 1.5	1.1	1690–2530	1959 \pm 162
PS/ α -CD	15	α -CD, 21.3%	66.9 \pm 0.3	2.2	200–2280	940 \pm 490
PS/ β -CD	15	β -CD, 25%	60.6 \pm 0.2	3.8	474–2840	1350 \pm 480
PS/ γ -CD	15	γ -CD, 28.5%	67.9 \pm 0.4	1.7	310–1830	967 \pm 380

^aWith respect to solvent (DMF). ^bWith respect to the polymer (PS).

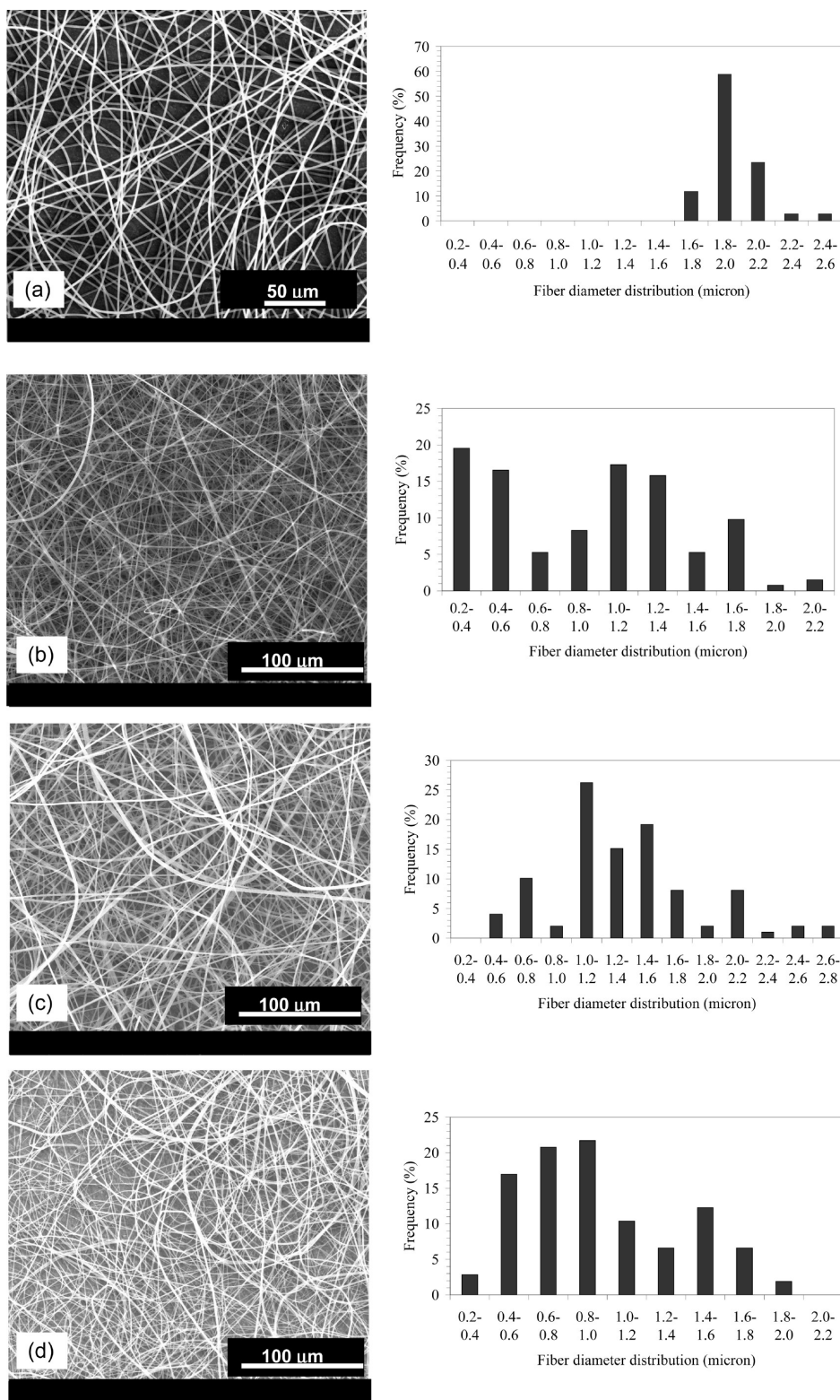


Figure 2. Representative SEM images of (a) PS, (b) PS/α-CD, (c) PS/β-CD, and (d) PS/γ-CD electrospun fibers. The fiber diameter distributions are shown on the right side.

rized in Table 1. The representative scanning electron microscopy (SEM) images of the electrospun PS and PS/CD fibers and the fiber diameter distributions are shown in Figure 2. It was observed that the diameter of the resulting electrospun PS/CD fibers is more or

less close to each other without showing any considerable differences except the PS/β-CD fibers are slightly thicker compared to PS/α-CD and PS/γ-CD fibers. In the case of PS fibers without CD, the fibers were thicker compared to PS/CD fibers since higher polymer concen-

TABLE 2. Atomic Concentrations Generated from XPS Wide Energy Survey Scans

Sample	C/%	O/%
PS	100	0
α -CD	59.6	40.4
β -CD	58.7	41.3
γ -CD	58.4	41.6
PS/ α -CD nanoweb	98.3	1.7
PS/ β -CD nanowebs	98.0	2.0
PS/ γ -CD nanoweb	97.8	2.2

tration was used in order to obtain bead-free fibers.²⁶ In brief, the variations in fiber diameters are due to differences in conductivity and viscosity of the polymer solutions.²⁶

XPS and ToF-SIMS Characterization. The presence of CD molecules at the surface of the fibers has a large influence on the molecular filtration capability. XPS was used to quantitatively obtain information of the surface composition of the PS/CD nanofibers, and ToF-SIMS was used to provide molecular and spatial distribution information on individual fiber surfaces. Table 2 shows elemental compositions based on wide energy survey spectra of the surface of PS, pure CDs, and the three different electrospun PS/CD nanoweb samples. The presence of low percentages of oxygen on the electrospun PS/CD nanofibers thus confirms the localization of the CD within the outer surface layers.

To provide more chemical state information of the surface chemistry, high-resolution C 1s spectra were recorded for each sample. Figure 3 shows C 1s spectra of PS, α -CD, and PS/ α -CD. The spectra are identical to those acquired for PS/ β -CD and PS/ γ -CD samples (data not shown). The C 1s spectrum of pure PS (Figure 3C) can be fitted with two peaks at binding energies at 284.7 and 285.0 eV and a smaller peak at 291.4 eV (marked I). The latter is the shakeup satellite arising from $\pi \rightarrow \pi^*$ transitions in the aromatic rings of the PS molecule. The two components for the main peak are assigned to the aromatic carbon $C=C-C=C$ at 284.7 eV (marked V) and the aliphatic carbon at 285.0 eV (marked IV). The spectrum of pure CD (Figure 3B) is fitted with three main components: the aliphatic carbon at 285.0 eV (small hydrocarbon contamination marked IV), $C-O-C/OH$ at 286.8 eV (marked III), and $O-C-O/C=O$ at 288.1 eV (marked II). The PS/ α -CD fiber surface can be successfully described as a combination of the spectra for the pure materials where the presence of CD is seen as a small shoulder (marked III) on the higher binding energy side of the main, PS carbon peak (Figure 3A). Thus, it is assumed that the elemental compositions generated from wide energy survey spectra are linear combinations of the elemental compositions of the materials. On the basis of this assumption, it is estimated that the approximately 5% of the probed surface consists of CD molecules for all three

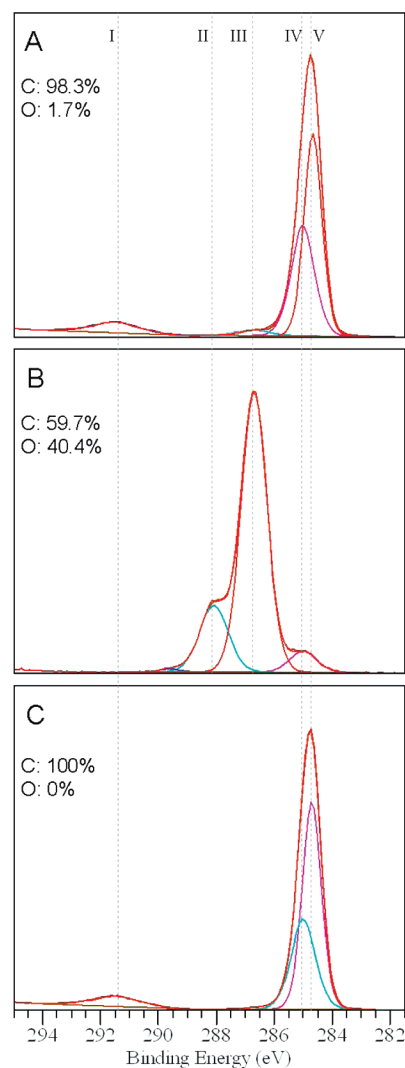


Figure 3. C 1s XPS spectra of (A) PS/ α -CD, (B) α -CD, and (C) PS.

different PS/CD samples. A slight increase in surface content (from 4.2 to 5.3%) is observed when going from α -CD through β -CD to γ -CD. The increase is correlated with the mass of CD used since samples were prepared from solutions with the same molar concentration of CD, which corresponds to a higher mass content of γ -CD compared to α -CD. On the other hand, the surface content of CD for all three PS/CD samples is lower than the CD content of the solutions they were prepared from. This indicates that the CD molecules are not homogeneously mixed throughout the fiber matrix, and some CD molecules are buried in the bulk of the fibers. CD molecules could phase separate from PS matrix and formed heterogeneous dispersion during solvent evaporation in the electrospinning process. In fact, we prepared film from PS/CD solutions by spin coating and solvent casting method, but CD molecules were phase separated and formed aggregates immediately during the solvent evaporation. This is likely because CD has a hydrophilic characteristic and PS is a hydrophobic polymer.

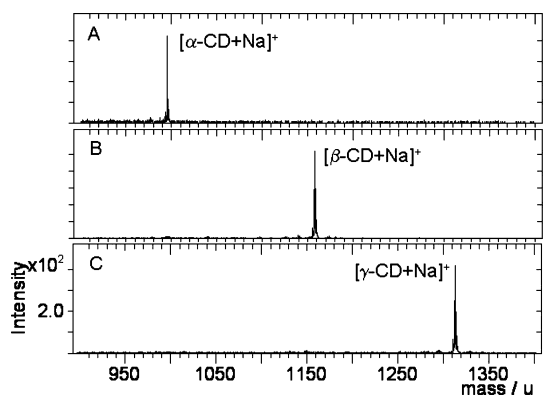


Figure 4. Positive ion *static*-ToF-SIMS spectra from PS/CD fiber samples in the mass range of 900–1400 units showing molecular CD peaks of (A) α -CD, (B) β -CD, and (C) γ -CD.

ToF-SIMS was used to corroborate the XPS results and to obtain information about the lateral distribution of CD in the outer molecular layer of fibers by chemical imaging. In addition, the probe depth of ToF-SIMS is 1–2 nm compared to 10 nm of XPS and thus is more surface-sensitive. Figure 4 shows positive ion ToF-SIMS mass spectra in the m/z range from 900 to 1400 units for each of the three PS/CD fiber samples. The

presence of CD is clearly established by peaks that can be assigned because of the high mass resolution achievable. In this case, the assignments can be attributed to the Na adduct ions of intact molecules of each of the CDs at atomic mass unit values of 995.24, 1157.46, and 1319.51 for α -CD, β -CD, and γ -CD, respectively.

High spatial resolution chemical imaging was performed on the fiber surfaces for PS/CD fiber samples. Since α -CD, β -CD, and γ -CD are found to have similar fragmentation patterns, the same peaks can be used for imaging of each sample. In order to enhance the chemical contrast of the images, the intensities of a number of fragment ions from either PS or CD are summed ($C_6H_5^+$, $C_7H_7^+$, $C_9H_9^+$, and $C_{15}H_{13}^+$ for PS; and $C_2H_5O_2^+$, $C_4H_5O^+$, and $C_3H_5O_2^+$ for CD). Thus two separate chemical images are created: one showing the location of PS and one showing CD, on the PS/CD fiber surface. Unfortunately, it was not possible to get sufficient contrast for the individual $[M + Na]^+$ peaks for each CD since the ion intensities were too low at the chosen raster size, which was necessary to see individual fibers and remain below the *static* limit (below the damage threshold).

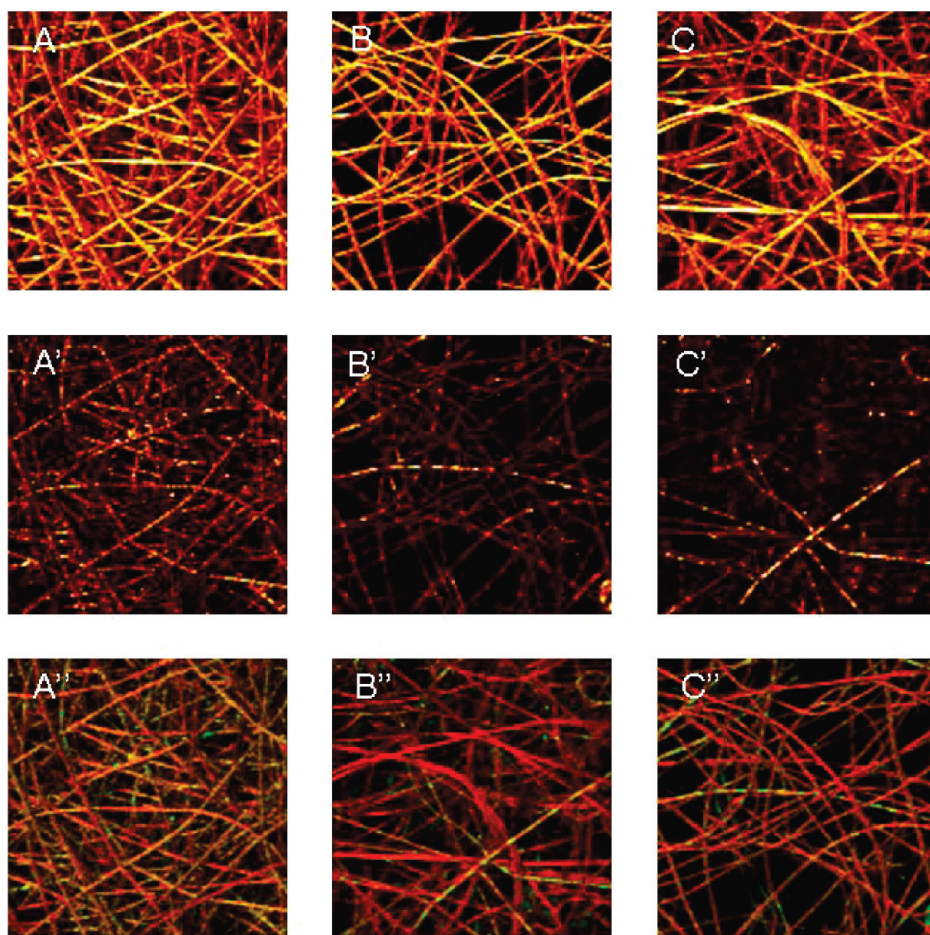


Figure 5. ToF-SIMS chemical images of the PS/CD fibers showing PS and CD. (A) PS fragment ion image of PS/ α -CD. (A') CD fragment ion image of PS/ α -CD. (A'') Overlay of the CD image (A', green) on the PS image (A, red) visualizing the location of CD on the PS fibers. B, B', and B'' show equivalent images for PS/ β -CD. C, C', and C'' show equivalent images for PS/ γ -CD. All images are $200 \times 200 \mu m^2$.

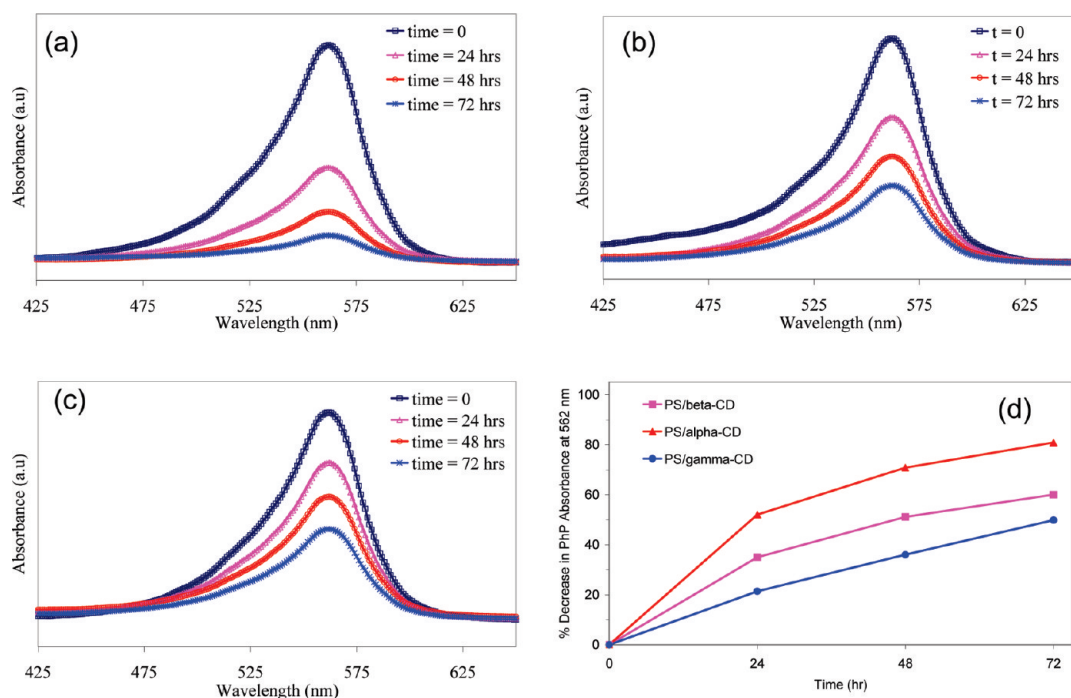


Figure 6. UV–vis spectra of phenolphthalein (Php) solution as a function of time after immersion of the nanoweb in (a) PS/α-CD, (b) PS/β-CD, (c) PS/γ-CD. (d) Cumulative % decrease of Php absorbance over time at 562 nm.

Figure 5 shows PS and CD chemical images for the three PS/CD fiber samples. By comparing the CD image with the PS image for each sample and overlaying the images, the distribution of CD on the surface of the fibers can be visualized. It is seen that small adjacent regions of single fibers that look similar in the PS images appear with different contrasts in the CD images. The uneven distribution of the CD signal suggests that the CD molecules form clusters on the PS fiber surfaces. Furthermore, there are larger areas of individual fibers that are clearly enriched in PS but deficient in CD fragment ions. This uneven distribution of CD on the fibers is independent of CD type. As discussed previously, CD molecules have a heterogeneous dispersion throughout the PS fiber matrix due to its hydrophilic nature and phase separation from the hydrophobic matrix during the solvent evaporation in the electrospinning process.

The surface analysis results clearly indicate that there is a surface enrichment or segregation of CD molecules on the fibers. This effect has been observed before for electrospun fibers made from blends, where the low molecular weight (MWt) additive is surface-concentrated.²⁹ In addition, mixtures of polymer blends of different MWt have been shown to demix at the surface where the low MWt weight component enriches the surface due to energetic and entropy effects.^{30,31} In such systems, it is proposed that the lower energy components migrate to the surface and minimize the surface tension, and the process is enhanced when the low MWt component is branched or immiscible with the higher M_w component.^{30–32}

It is also noteworthy saying that CDs are physically incorporated in the fiber matrix; the chemical reaction between CD and PS is not expected. In addition, no evidence of chemical bonding of CD to PS matrix was observed from XPS, ToF-SIMS, and DP-MS analyses. Moreover, the inclusion complexation between CD and PS chains is excluded since the cavity of all three CDs (α-CD, β-CD, and γ-CD) is too narrow to encapsulate amorphous PS chains.^{33,34} So, the physically bounded CD molecules on the fiber surface are empty and available to perform their filtration function by capturing molecules from the surroundings. This is indeed one of the reasons for choosing PS as the fiber matrix because the matrix would not interfere in the complexation of CD with the target molecules to be trapped.

UV–Vis Measurements: Uptake of Phenolphthalein. The molecular filtration capability of PS/CD fibrous nanoweb was tested using phenolphthalein (Php) solutions as a model system because Php is known to form inclusion complexes with CDs and it is easy to follow its uptake by UV–vis measurements.^{35–38} The PS/CD fibrous webs were immersed into Php solution, and the change in absorbance of Php was recorded as a function of time by UV–vis spectrophotometry (Figure 6). It was observed that the absorbance of Php solution decreased significantly over time in the presence of PS/CD nanoweb due to the removal of Php from solution by the complexation with the CDs present on the fiber surface. When the CD types were compared (Figure 6a–c), the percentage removal of Php with respect to time was on the order of PS/α-CD > PS/β-CD > PS/γ-CD. The ToF-SIMS chemical mapping of PS/CD nanoweb shows that

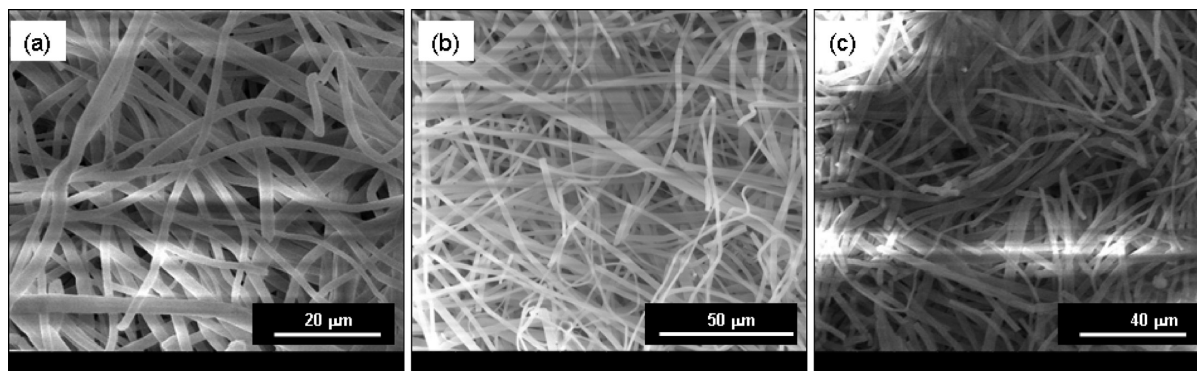


Figure 7. Representative SEM images of (a) PS/α-CD, (b) PS/β-CD, and (c) PS/γ-CD nanoweb after 3 days of immersion in phenolphthalein (Php) solution.

α-CD molecules have a more homogeneous distribution on the nanoweb surface compared to β-CD and γ-CD (Figure 5), and this may lead to the removal of Php more rapidly in the case of PS/α-CD nanoweb. Figure 6d summarizes the cumulative % decrease of Php ab-

sorbance over time. All three PS/CD nanoweb demonstrate the ability to function as a molecular filter through complexation of the Php molecules with the CD molecules. In our previous study, we have shown that the electrospun PS nanoweb without having CDs

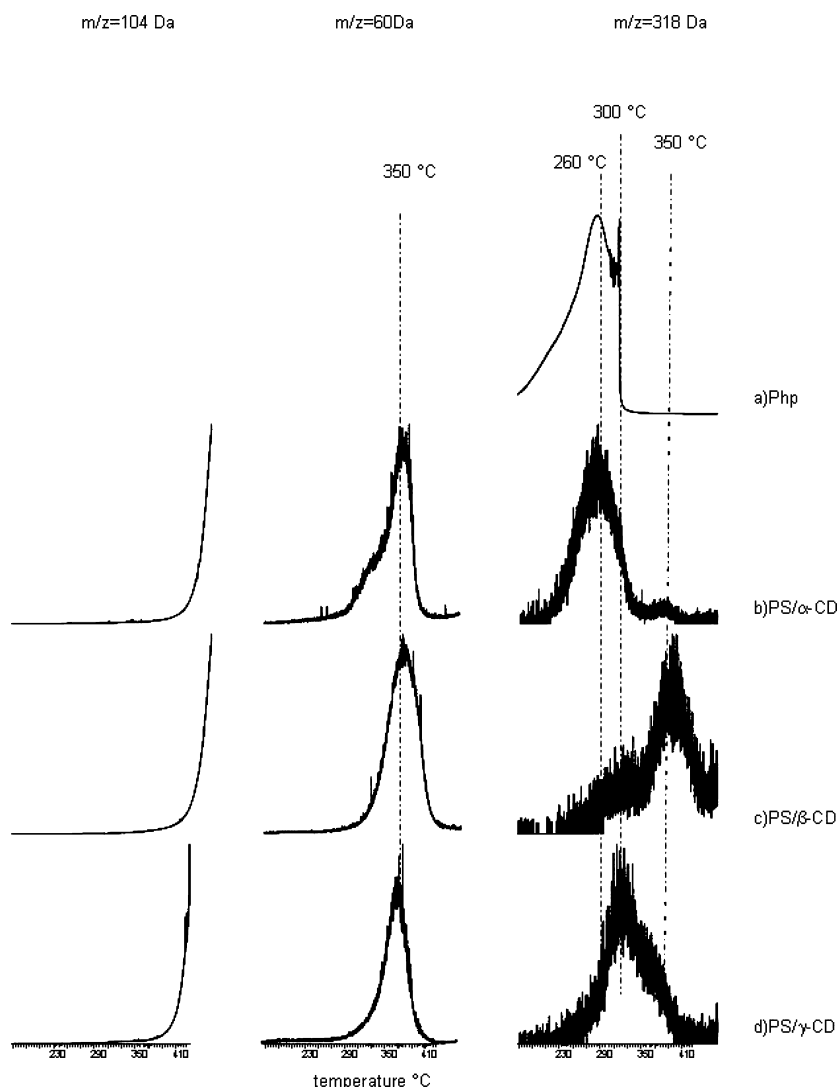


Figure 8. DP-MS evolution profiles of PS-based product; monomer ($m/z = 104$ Da), CD-based product; $C_2H_4O_2$ ($m/z = 60$ Da) and phenolphthalein (Php) ($m/z = 318$ Da), detected during the pyrolysis of (a) pure phenolphthalein, (b) PS/α-CD, (c) PS/β-CD, (d) PS/γ-CD. (Note: The fibers were analyzed after the exposure to Php solution at the end of the UV-vis experiments.)

were ineffective at trapping the Php from the solution.¹⁴ On the other hand, for the PS/CD nanowebs, the trapping of the Php was very significant, suggesting that these electrospun nanowebs would be very effective for filtering the molecules from the solutions. We have also examined the dimension stability of these PS/CD nanowebs, and we observed that the nanowebs kept their fibrous structure after the filtration process (Figure 7).

Direct Pyrolysis Mass Spectrometry. Direct pyrolysis mass spectrometry (DP-MS) studies were performed on the PS/CD nanowebs exposed to phenolphthalein (Php) solution to confirm the complexation of CDs with Php molecules. The thermal evaporation of the guest molecules shifts to higher temperatures due to the interactions with the CD cavity, thus, DP-MS is a useful technique to characterize the CD host–guest inclusion complexes.³⁹ In general, for a multicomponent system, DP-MS allows separation of components as a function of their volatilities and/or thermal stabilities.^{40–42} Thus, in pyrolysis MS analysis, not only the detection of a peak but also the variation of its intensity as a function of temperature and the evolution profile is important.^{39–42}

In Figure 8, evolution profiles of PS-, CD-, and phenolphthalein (Php)-based products, namely, the styrene monomer ($m/z = 104$ Da) of PS, $C_2H_4O_2$ ($m/z = 60$) for α -CD, β -CD, and γ -CD, and Php (molecular peak at $m/z = 318$ Da) recorded during the DP-MS analyses are given. For all of the samples, PS- and CD-based products showed identical behaviors with those of the corresponding pure forms (Figure 8). On the other hand, the trends observed in the evolution profiles of Php-based products showed noticeable differences for PS/CD samples. For the PS/ α -CD, the evolution profile of Php was almost identical to that of the pure Php with a maximum at 260 °C, except the presence of a second and a weak peak at around 350 °C (Figure 8b). In the case of the sample containing β -CD, two peaks with maxima at around 295 and 350 °C, the first being weaker, were observed (Figure 8c). For the sample containing γ -CD, a broad peak with a maximum at around 295 °C and a shoulder at around 350 °C were recorded (Figure 8d). It is clear that the weakest interaction was the one between Php and α -CD; for this sample, the

evolution of most of the Php occurred independently as in case of the pure form. On the other hand, for both of the samples involving β -CD or γ -CD, the evolution of Php shifted to higher temperatures and the trends in the evolution profiles indicated the presence of two different environment/interactions of Php with CD cavities. When the trends in the evolution profiles are compared, it can be concluded that the interaction between Php molecule and CD cavity is on the order α -CD < γ -CD < β -CD, indicating that the size of the CD cavity and the size of the host molecule were critical for the strength of an inclusion complex.^{21,22} This finding suggests that the binding between the CD cavity and the phenolphthalein is the strongest for β -CD, and the reason for that is the proper size/shape match between β -CD and phenolphthalein, which correlates with the literature findings.^{35–37}

CONCLUSION

A comparative study has been carried out on the molecular filtration capabilities of electrospun polystyrene nanofibers functionalized with α -CD, β -CD, and γ -CD molecules. The capability of the PS/CD nanofibers/nanowebs to filter molecules from solution was determined by using phenolphthalein as a model compound. Chemical surface analysis using XPS and static-ToF-SIMS demonstrated that the CD molecules were present at the surface of the PS fibers. The relative binding strengths of the different CDs for phenolphthalein within the PS fiber matrix were determined using direct pyrolysis mass spectrometry (DP-MS). The results indicate that there is a trend depending on the CD cavity size, and the strength of interaction between phenolphthalein and the CD cavity is on the order of α -CD < γ -CD < β -CD. Such information is invaluable for designing cyclodextrin functionalized electrospun nanowebs where the properties of the fiber surfaces and inclusion complex formation are crucial for optimizing the efficiency of the filtration performance. In conclusion, our findings are very promising and show the potential application for cyclodextrin containing electrospun nanofibers/nanowebs to be used as molecular filters for removal of organic molecules from solutions.

METHODS

Materials. Polystyrene (PS) ($M_w \sim 280\,000$), *N,N*-dimethylformamide (DMF) (99%), phenolphthalein (ACS reagent), and ethanol (absolute, HPLC grade, $\geq 99.8\%$) were purchased from Sigma-Aldrich. The α -, β -, and γ -cyclodextrins (α -CD, β -CD, and γ -CD) were obtained from Wacker Chemie AG (Germany). The materials were used as received.

Electrospinning. PS/CD solutions were prepared by dissolving PS and CDs (α -CD, β -CD, and γ -CD) in DMF. The polymer concentration was 15% w/v (with respect to solvent), and the CD content was 21.3, 25, and 28.5% (w/w) (with respect to polymer) for α -CD, β -CD, and γ -CD, respectively. Different weight ratios were used for CDs in order to keep the

molar ratio the same in the polymer matrix. In the case of PS without CD, 25% w/v polymer solution was used in order to get bead-free uniform fibers. The PS and PS/CD solutions were placed in a syringe fitted with a metallic needle (0.4 mm inner diameter). The syringe was fixed horizontally on the syringe pump (model KDS 101, KD Scientific), and the high voltage power supply (Spellman, MP Series) was used. The applied electrospinning parameters were as follows: applied voltage = 15 kV, feed rate = 1 mL/h, and tip-to-collector distance = 10 cm. A grounded stationary rectangular metal collector plate covered by a piece of aluminum foil was used for the fiber deposition. The collected fibers were dried at 40 °C under vacuum oven for 24 h to remove the residual solvent if present.

Characterization. The viscosity of the solutions was measured at 24 °C using the Brookfield DV-III Ultra rheometer, which is equipped with a cone/plate accessory of spindle type CPE-41. The viscosity measurements were repeated three times to check the reproducibility and the consistency of the viscosity reading. The conductivity of the solutions was measured with multiparameter meter InoLabMulti 720 (WTW) at room temperature.

The morphologies of the cyclodextrin functionalized electrospun polystyrene (PS/CD) fibers were investigated by scanning electron microscopy (SEM) (FEI, Nova 600 NanoSEM and Quanta 200 FEG). The fiber diameter distribution was determined from the SEM images, and around 100 fibers were analyzed.

X-ray photoelectron spectroscopy (XPS) measurements were carried out using a Kratos Axis Ultra^{DL}. Samples for XPS were prepared by electrospinning a covering mat of fibers onto Al foil, and the samples were analyzed on the foil. A monochromated Al K α X-ray source was used ($h\nu = 1486.6$ eV) and operated at 10 mA and 15 kV. A hybrid lens mode was employed during analysis (electrostatic and magnetic), with an analysis area of approximately $300\ \mu\text{m} \times 700\ \mu\text{m}$. The analysis takeoff angle (TOA) was 0° (with respect to the Al foil surface), allowing a maximum probe depth (10 nm). Wide energy survey scans were obtained over the range of 0–1200 eV binding energy (BE) at a pass energy of 160 eV and used to determine the surface elemental composition. High-resolution spectra were recorded for carbon 1s at a detector pass energy of 20 eV. The Kratos charge neutralizer system was used on all samples with a filament current between 1.8 and 2.1 A and a charge balance of 3.6 V.

ToF-SIMS was performed using an ION-TOF TOF.SIMS V instrument equipped with a Bi primary ion cluster source operating at 25 kV. Bi $_3^{2+}$ primary ions were used during analysis using the high current bunched mode for maximum mass resolution. Samples for ToF-SIMS were prepared by electrospinning a few layers of fibers directly onto pieces of Si wafers. Chemical imaging was performed with the burst alignment mode, which offers a high spatial resolution but low mass resolution. Chemical imaging over an area of $200 \times 200\ \mu\text{m}^2$ was based only on peaks unambiguously identified in previous high mass resolution measurements.¹⁴ A low energy electron flood gun was used for charge compensation, and the primary ion dose was kept below 10^{12} ions/cm 2 to stay below the static limit.

The molecular filtration capability of the PS/CD electrospun fibrous nanowebs was tested by using phenolphthalein (Php) as a model organic molecule. The uptake of Php by PS/CD nanowebs was determined by measuring the reductions in absorbance from solution using UV–vis spectrophotometry (Helios- β). In brief, 4×10^{-4} M of Php solution was prepared in absolute ethanol, and the pH of the solution was adjusted to pH 11 by addition of buffer solution. About 16 mg of PS/CD fibers was placed separately in the bottom of the UV–vis cuvettes filled with purple color Php solution. The UV cuvettes were covered tightly with a Teflon lid to prevent the evaporation of solution, and they were not disturbed for 3 days. The absorbance spectra of Php solution were recorded initially (time = 0, right after dipping the fibers in the cuvette) and after every 24 h. The PS/ β -CD nanoweb was also tested for 6 days, and it was observed that the decrease in PhP absorption was about the same after 3 days and after 6 days, indicating that the saturation point was reached after approximately 3 days; therefore, the immersion experiments were stopped at the end of 3 days. After 3 days, the fibrous mats were removed, rinsed with ethanol and water to remove physically adsorbed PhP molecules, and then analyzed by direct pyrolysis mass spectrometry (DP-MS). The pH of the solutions was also measured before and after the UV–vis experiments, and it was observed that the pH of the solution was unchanged at the end of UV–vis experiments.

The DP-MS system consists of Waters Quattro Micro GC tandem MS with an EI ion source and a mass range of 10–1500 Da coupled with a direct insertion probe ($T_{\text{max}} = 650$ °C); 0.01 mg of each fiber sample was pyrolyzed in flared quartz sample vials. The temperature was increased at a rate of 10 °C/min, and the scan rate was 1 scans/s, with simultaneous mass spectrometric analysis of the pyrolytic fragments.

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